In the claims:

For the convenience of the Examiner, all claims being examined are presented below. Please cancel claim 2 without prejudice.

- 1. **(Amended)** A method for identifying an inhibitor compound capable of reducing the interaction between a first region and a second region, comprising:
- a) placing in contact:
 - i) a potential inhibitor compound;
- ii) a first region which is a fragment of a nuclear protein, wherein the fragment comprises a signature motif B'XXLL, in which B' is any natural hydrophobic amino acid, L is leucine, and X independently represents any natural amino acid, and the signature motif is a structural element of a nuclear protein that binds to a liganded nuclear receptor in the process of activating or repressing target genes, and the nuclear protein is a bridging factor responsible for an interaction between a liganded nuclear receptor transcription factor and a transcription initiation complex involved in regulation of gene expression; provided that a fragment that includes residues 624-1287 of TIF-2 is excluded;
- iii) a second region which is a liganded nuclear receptor transcription factor or a fragment thereof, wherein the fragment comprises that part of the nuclear receptor transcription factor which is capable of interacting with a nuclear protein through binding to the signature motif; and
- b) detecting the presence or absence of inhibition of the interaction between ii) and iii).
- 3. (Amended) A method according to claim 23, wherein B¹ is leucine or valine.
- 4. (Amended) A method according to claim 3, wherein B¹ is leucine.
- 5. (Amended) A method according to claim 1, 3, 4, or 23, wherein the signature motif is B²B¹XXLL, wherein B² is a hydrophobic amino acid.
- 6. (Amended) A method according to claim 5, wherein B² is selected from isoleucine, leucine, methionine, phenylalanine, tryptophan, tyrosine and valine.
- 7. **(Amended)** A method according to claim 1, 3, 4, or 23, wherein the nuclear protein is a coactivator.

- 8. (Amended) A method according to claim 7, wherein the coactivator is selected from RIP 140, SRC-1, TIF2, CBP, p300, TIF1, Trip1, Trip2, Trip3, Trip4, Trip5, Trip8, Trip9, p/CIP, ARA70 & Trip230.
- 9. (Amended) A method according to claim 1, 3, 4, or 23, wherein the transcription factor is a steroid hormone receptor.
- 10. **(Amended)** A method according to claim 9, wherein the steroid hormone receptor is selected from oestrogen receptor, progesterone receptor, androgen receptor and glucocorticoid receptor.
- 11. **(Amended)** A method according to claim 10, wherein the steroid hormone receptor is oestrogen receptor.
- 12. **(Amended)** A method according to claim 1, 3, 4, or 23, wherein the method is a 2-hybrid assay.
- 13 (Amended) A method according to claim 1, 3, 4, or 23, wherein the potential inhibitor compound is a member of a peptide library based on the signature motif.

Please add the following new claim:

23. (New) A method according to claim 1, wherein the fragment of a nuclear protein comprises only one signature motif.

The amended claims are restated below to reflect changes from the last filing.

- 1. (Amended) A method for identifying <u>an</u> inhibitor compounds-capable of reducing the interaction between a first region and a second region, comprising:
- a) placing in contact:
 - i) a potential inhibitor compound;
- ii) a first region which is a <u>fragment of signature motif on</u> a nuclear protein, <u>wherein</u> the <u>fragment comprises a signature motif B'XXLL</u>, in which B' is any natural hydrophobic amino acid, L is leucine, and X independently represents any natural amino acid, and the <u>signature motif is a structural element of a nuclear protein that binds to a liganded nuclear</u>

receptor in the process of activating or repressing target genes, and the nuclear protein is a bridging factor responsible for an interaction between a liganded nuclear receptor transcription factor and a transcription initiation complex involved in regulation of gene expression; provided that a fragment that includes residues 624-1287 of TIF-2 is excluded;

[b)] <u>iii)</u> a second region which is a liganded nuclear receptor transcription factor or a fragment thereof, wherein the fragment comprises that part of the a nuclear receptor which is capable of interacting with the nuclear protein through binding to the signature motif; [,] and wherein:

the nuclear protein is a bridging factor that is responsible for the interaction between a liganded nuclear receptor and a transcription initiation complex involved in regulation of gene expression; the nuclear receptor is a transcription factor;

the signature motif is a short sequence of amino acid residues which is the key structural element of a nuclear protein which binds to a liganded nuclear receptor as part of the process of the activation or repression of target genes; and

in which the method comprises taking:

- i) the potential inhibitor compound;
- ii) the liganded nuclear receptor or a fragment thereof in which the fragment comprises the second region defined in this claim in b) above;
- iii) a fragment comprising a signature motif of the nuclear protein; and
- b) detecting the presence or absence of inhibition of the interaction between ii) and iii).
- 3. (Amended) A method according to claim 23 2 or claim 5, wherein in which B' is leucine or valine.
- 4. (Amended) A method according to claim 3, wherein in which B¹ is leucine.
- 5. (Amended) A method according to claim 1, 3, 4, or 23 wherein in which the signature motif is B²B¹XXLL wherein B² is any natural hydrophobic amino acid, B² is a hydrophobic amino acid, L is leucine and X independently represents any natural amino acid.
- 6. (Amended) A method according to claim 5, wherein in which B² is selected from the group consisting of isoleucine, leucine, methionine, phenylalanine, tryptophan, tyrosine and valine.
- 7. (Amended) A method according to any one of claims 1, 3, 4, or 23 and 5, wherein in which the nuclear protein is a coactivator.